TECH CENTER 1507/2900

displaying a second axis substantially perpendicular to said first axis, said second axis indicating expression level in said second sample;

for a selected expressed sequence, displaying a mark at a position with an X coordinate and a Y coordinate, wherein the X coordinate of said position is selected relative to said first axis and said Y coordinate of said position is selected relative to said second axis, wherein said position is selected relative to said first axis in accordance with an expression level of said selected expressed sequence in said first sample and relative to said second axis in accordance with an expression level of said selected expressed sequence in said second sample;

receiving an input of a user's selection of said mark; and in response to said user input, displaying information about said selected expressed sequence.

- 2. The method of Waim 1 wherein said selected expressed sequence comprises a gene.
- The method/of claim 1 wherein said selected expressed sequence 3. comprises a portion of a gene.
- 4 (Previously Amended). The method of claim 1 further comprising: repeatedly displaying a mark for each one of a plurality of selected expressed sequences.
- 5 (Previously Amended). The method of claim 1 further comprising: monitoring said expression level of said expressed sequence in said first sample and said second sample.
- 6 (Previously Amended). The method of claim 5 wherein said monitoring further comprises

inputting a plurality of hybridization intensities from pairs of perfect match and mismatch probes, said perfect match probes being perfectly complementary to a target nucleic acid sequence/indicative of expression of said selected gene and said mismatch probes having

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David H. Mack Application No.: 09/020,743 Page 3 at least one base mismatch with said target sequence, and said hybridization intensities 6 7 indicating hybridization affinity between said perfect match and mismatch probes and a sample 8 nucleic acid sequence from said one of said samples; 9 comparing the hybridization intensities of each pair of perfect match probe and 10 mismatch probe; and 11 generating said expression level for said expressed sequence and said one of said samples responsive to results of said comparing. 12 1 7 (Previously Amended). The method of claim 6 further comprising: 2 comparing a difference between hybridization intensities of perfect match and 3 mismatch probes at a base position to a difference threshold. 1 8 (Previously Amended). The method of claim 7 further comprising: 2 comparing a quotient of hybridization intensities of perfect match and mismatch 3 probes at a base position to a ratio threshold. 1 9 (Previously Amended). The method of claim 6 further comprising: 2 counting a probe pair as a positive probe pair to increment a positive probe pair count if a perfect match probe intensity minus a mismatch probe intensity exceeds a 3 4 difference threshold and said perfect match probe intensity divided by said mismatch probe 5 intensity exceeds a ratio threshold; 6 counting said probe pair as a negative probe pair to increment a b) 7 negative probe pair count If said mismatch probe intensity minus said perfect match probe intensity exceeds said difference threshold and said mismatch probe intensity divided by said 8 9 perfect match probe intensity exceeds said ratio threshold; 10 computing a logarithmic ratio of said perfect match probe intensity to c) 11 said mismatch probe Intensity; and 12 d) computing a difference of said perfect match probe intensity to said 13 mismatch probe intensity. 10/(Previously Amended). 1 The method of claim 9 further comprising:

David H. Mack Application No.: 09/020,743 Page 4 2 3 exhibit said difference; and 4 5 6 average of said differences. 1 12 (Previously Amended). 1 2 3 1 13 (Previously Amended). 2 14. 1 2 3 sequence. 15. 1 2 1 16 (Previously Amended). 2 characteristic comprises a disease state. 17. 1 2 a treatment strategy for a disease. 1 18. 2 a disease.

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repeating said a), b), c) and d) steps for each of said probe pairs, accumulating a sum of differences of said perfect match and mismatch probe intensities for probe pairs that

determining an expression level of said selected expressed sequence to be an

11. This claim has been previously canceled.

The method of claim 1 further comprising: in response to said input, displaying information about said selected expressed sequence; said information comprising an identifier for said selected expressed sequence.

The method of claim 12 wherein said identifier for said selected expressed sequence comprises a GenBank accession number.

- The method of claim 12 wherein said information about said selected expressed sequence comprises a/GenBank database record for said selected expressed
- The method of claim 1 wherein said first sample and said second sample are collected from tissue samples differing in a particular characteristic.

The method of claim 15 wherein said particular

- The method of claim 15 wherein said particular characteristic comprises
- The method of claim 1 wherein said particular characteristic is a stage of

19 (Previously Amended). The method of claim 1 further comprising: displaying a third axis substantially perpendicular to said first axis and to said second axis in a three-dimensional display environment wherein said position of said mark is

David H. Mack Application No.: 09/020,743 Page 5 further selected relative to said third axis in accordance with an expression level of said 4 5 selected expressed sequence in a third sample. A computer-implemented method of presenting 20 (Previously Amended). 1 2 sample analysis information comprising: displaying a first axis indicating a concentration of a compound in a first sample 3 as determined by monitoring binding of said compound to a selected polymer having binding 4 5 affinity to said compound; displaying a second axis substantially perpendicular to said first axis, said 6 7 second axis indicating a concentration of said compound in said second sample as determined 8 by monitoring binding of said compound to said selected polymer; and 9 displaying a mark at a position with an X coordinate and a Y coordinate, wherein the X coordinate of said position is selected relative to said first axis and said Y 10 11 coordinate of said position is selected relative to said second axis, wherein said position is 12 selected relative to said first axis in accordance with said concentration in said first sample and 13 relative to said second axis in accordance with said concentration in said second sample; 14 receiving an input of a user's selection of said mark; and 15 in response to said user input, displaying information about said selected 16 expressed sequence. 21. 1 The method of claim 20 wherein said selected polymer comprises a 2 nucleic acid sequence. 22. 1 The method of claim 20 wherein said selected polymer comprises a 2 protein. 1 23 (Previously Amended). The method of claim 21 further comprising: 2 obtaining said concentration of said compound in said first sample by exposing 3 said first sample to a plurality of nucleic acid probes. 1 24 (Previously Amended). The method of claim 22 further comprising:

David H. Mack Application No.: 09/020,743 Page 6 obtaining said concentration of said compound in said first sample by exposing 2 3 said first sample to a plurality of peptide probes. 1 25 (Previously Amended). A computer program product for presenting expression level information as collected from a first sample and a second sample, said product 2 3 comprising: code for displaying a first axis indicating expression level in said first sample; 4 code for displaying a second axis substantially perpendicular to said first axis, 5 6 said second axis indicating expression level in said second sample; 7 code for, for a selected expressed sequence, displaying a mark at a position with 8 an X coordinate and a Y coordinate, wherein the X coordinate of said position is selected 9 relative to said first axis and said Y coordinate of said position is selected relative to said 10 second axis, wherein said position is selected relative to said first axis in accordance with an 11 expression level of said selected expressed sequence in said first sample and relative to said 12 second axis in accordance with an expression level of said selected expressed sequence in said MIGHTS 13 second sample; 14 code for receiving an input from a user's selection of said mark; code for displaying information about said selected expressed sequence in 15 16 response to said user input; and 17 a computer-readable storage medium for storing the codes. 1 26. The product of claim 25 wherein said selected expressed sequence 2 comprises a gene. The product of claim 25 wherein said selected expressed sequence 27. 1 2 comprises a portion of a gene. 1 28. The product of claim 25 further comprising code for repeatedly applying 2 said displaying a mark code for a plurality of selected expressed sequences. 29. The product of claim 25 further comprising: 1

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code for monitoring said expression level of said expressed sequence in said first sample and said second sample.

The product of claim 29 wherein said code for 30 (Previously Amended). monitoring for one of said samples comprises:

code for inputting a plurality of hybridization intensities from pairs of perfect match and mismatch probes, said perfect match probes being perfectly complementary to a target nucleic acid sequence indicative of expression of said selected gene and said mismatch probes having at least one base mismatch with said target sequence, and said hybridization intensities indicating hybridization affinity between said perfect match and mismatch probes and a sample nucleic acid sequence from said one of said samples;

code for comparing the hybridization intensities of each pair of perfect match probe and mismatch probe; and

code for generating said expression level for said expressed sequence and said one of said samples responsive to a result produced by said code for comparing [step].

> The product of claim 30 further comprising: 31.

code for comparing/a difference between hybridization intensities of perfect match and mismatch probes at a base position to a difference threshold.

> 32. The product of claim 31 further comprising:

code for comparing a quotient of hybridization intensities of perfect match and mismatch probes at a base position to a ratio threshold.

- 33. The product of claim 30 further comprising:
- a) code for counting a probe pair as a positive probe pair to increment a positive probe pair count if a perfect match probe intensity minus a mismatch probe intensity exceeds a difference threshold and said perfect match probe intensity divided by said mismatch probe intensity exceeds a ratio threshold;
- b) code for counting said probe pair as a negative probe pair to increment a negative probe pair count if said mismatch probe intensity minus said perfect match probe

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intensity exceeds said difference threshold and said mismatch probe intensity divided by said perfect match probe intensity exceeds said ratio threshold; and

c) code for computing a logarithmic ratio of said perfect match probe intensity to said mismatch probe intensity.

34 (Previously Amended). The product of claim 33 further comprising: code for repeatedly applying said a), b), and c) codes for each of said probe pairs, accumulating a sum of differences of said perfect match and mismatch probe intensities for probe pairs that exhibit said difference; and

code for determining an expression level of said selected expressed sequence to be an average of said differences.

35. This claim has been previously canceled.

36 (Previously Amended). The product of claim 25 further comprising: code for, in response to said input, displaying information about said selected expressed sequence; said information comprising an identifier for said selected expressed sequence.

37 (Previously Amended). The product of claim 36 wherein said identifier for said selected expressed sequence comprises a GenBank accession number.

- 38. The product of claim 36 wherein said information about said selected expressed sequence comprises a GenBank database record for said selected expressed sequence.
- 39. The product of claim 25 wherein said first sample and said second sample are collected from tissue samples differing in a particular characteristic.

40 (Previously Amended). The product of claim 39 wherein said particular characteristic comprises a disease state.

The product of claim 39 wherein said particular characteristic comprises a treatment strategy for a disease.

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42 (Previously Amended). The product of claim 39 wherein said particular characteristic is a stage of a disease.

43 (Previously Amended). The product of claim 25 further comprising: code for displaying a third axis substantially perpendicular to said first axis and to said second axis in a three-dimensional display environment wherein said position of said mark is further selected relative to said third axis in accordance with an expression level of said selected expressed sequence in a third sample.

44. A computer program product for presenting sample analysis information comprising:

code for displaying a first axis indicating a concentration of a compound in a first sample as determined by monitoring binding of said compound to a selected polymer having bonding affinity to said compound;

code for displaying a second axis substantially perpendicular to said first axis, said second axis indicating a concentration of said compound in a second sample as determined by monitoring binding of said compound to said selected polymer;

code for displaying a mark at a position with an X coordinate and a Y coordinate, wherein the X coordinate of said position is selected relative to said first axis and said Y coordinate of said position is selected relative to said second axis, wherein said position is selected relative to said first axis in accordance with said concentration in said first sample and relative to said second axis in accordance with said concentration in said second sample;

code for receiving an input of a user's selection of said mark;

code for displaying information about said selected expressed sequence in response to said user input, and

a computer-readable storage medium that stores the codes.

45. The product of claim 44 wherein said selected polymer comprises a nucleic acid sequence.

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46. The product of claim 44 wherein said selected polymer comprises a protein.

47 (Previously Amended). A computer system comprising a display, a processor, and a memory that stores instructions for configuring said processor to:

display a first axis indicating expression level in said first sample;

display a second axis substantially perpendicular to said first axis, said second axis indicating expression level in said second sample; and

for a selected expressed sequence, display a mark at a position with an X coordinate and a Y coordinate, wherein the X coordinate of said position is selected relative to said first axis and said Y coordinate of said position is selected relative to said second axis, wherein said position is selected relative to said first axis in accordance with an expression level of said selected expressed sequence in said first sample and relative to said second axis in accordance with an expression level of said selected expressed sequence in said second sample; wherein information about said selected expressed sequence is displayed responsive to an input of a user's selection of said mark.

48 (Previously Amended). A computer system comprising a display, a processor, and a memory that stores instructions for configuring said processor to:

display a first axis indicating a concentration of a compound in a first sample as determined by monitoring binding of said compound to a selected polymer having binding affinity to said compound;

display a second axis substantially perpendicular to said first axis, said second axis indicating a concentration of said compound in said second sample as determined by monitoring binding of said compound to said selected polymer; and

display a mark at a position with an X coordinate and a Y coordinate, wherein the X coordinate of said position is selected relative to said first axis and said Y coordinate of said position is selected relative to said second axis, wherein said position is selected relative to said first axis in accordance with said concentration in said first sample and relative to said second axis in accordance with said concentration in said second sample; wherein information

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about said selected expressed sequence is displayed responsive to an input of a user's selection of said mark.

49. The method of claim 1 further comprising:

providing a tactile feedback to said user through a pointing device when a cursor is moved over said mark; said tactile feedback indicating expression level for said selected expressed sequence corresponding to said mark.

50. The method of claim / further comprising:

expressed sequence corresponding to said mark.

providing an aural indication to said user through a pointing device when a cursor is moved over said mark; said aural indication indicating expression level for said selected expressed sequence corresponding to said mark.

51. The method of claim 1 further comprising: obtaining information from an internet based resource about a selected

52. The method of claim 1 further comprising:
receiving from the user a selection of at least two of a plurality of marks, said
marks;

displaying information about genes corresponding to said selection of at least two of a plurality of marks.

REMARKS

The pending claims are 1-10, 12-34, and 36-52.

The Examiner rejected the pending claims under 35 U.S.C. § 103(a) as being unpatentable over Zhao et. al. (Gene Vol. 156 pp. 207-213 1995) in view of the various references to Seilhamer et al. (U.S. Pat. No. 6,023,659), Lockhart et al. (WO97/27317), Beattie et al. (U.S. Pat. No. 5,843,767), and Rosenberg et al. (U.S. Pat. No. 6,028,593).

Zhao discusses a method for analyzing large numbers of cDNA plasmids from brain tissue. The Zhao method includes preparing a filter using cDNA plasmids isolated from

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